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(54) **METHOD FOR MANUFACTURING A COMPOSITE SORBER FOR THE REMOVAL OF H2O CONSISTING OF HYGROSCOPIC INORGANIC SALTS DISSOLVED IN A POLYMERIC MATRIX**

VERFAHREN ZUR HERSTELLUNG EINES VERBUNDSORPTIONSMITTELS ZUR ENTFERNUNG VON H2O, BESTEHEND AUS HYGROSKOPISCHEN ANORGANISCHEN SALZEN, DIE IN EINER POLYMERMATRIX GELÖST SIND

PROCÉDÉ DE FABRICATION D'UN DÉSHYDRATEUR COMPOSITE POUR L'ÉLIMINATION D'H2O CONSTITUÉ DE SELS INORGANIQUES HYGROSCOPIQUES DISSOUS DANS UNE MATRICE POLYMÈRE

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Description

[0001] The presence of H₂O, even at trace levels, results to be harmful for the correct functioning of various devices, among which lithium batteries, microelectromechanical devices known in the field by the acronym MEMS (MicroElectroMechanical Systems), organic displays of the OLED-type (Organic Light Emitting Diode) and photovoltaic cells, i.e. the cells of the OSC-type (Organic Solar Cells), to mention some of the most interesting devices. In the following these devices, and more in general any sealed device in which the presence of H₂O results to be harmful, even at low levels (lower than 5000 ppm), are referred to by the term of sensitive device.

[0002] In the sensitive devices the presence of H₂O can cause the progressive deterioration of the performances; as an example, further information with respect to the effects of this contaminant can be found in the article "Correlation between dark spot growth and pinhole size in organic light-emitting diodes", by Shuang Fang Lim, et al., published in Applied Physics Letters, volume 78 no. 15, on 9 April 2001 as relating to the OLED displays, and in the fifth chapter of the book "Organic Photovoltaics - Concepts and Realization" by Brabec et al., edition of 2003 by Springer-Verlag, as relating to the photovoltaic cells of the OSC-type.

[0003] The use of sorbers for the removal of gaseous impurities from the housings of devices sensitive to their presence is known in the art. For example the international patent application WO 2004/072604 in the name of the applicant discloses the use of active components dispersed in suitable porous matrices; the patent applications WO 2007/013118 and WO 2007/013119, both in the name of the applicant, instead describe nanostructured systems in which the active component is confined in a porous medium, itself being dispersed in a polymeric matrix, whereas the use of functionalized nuclei dispersed in a permeable polymeric matrix is described in the international patent application WO 2007/074494 in the name of applicant.

[0004] All of the foregoing technological solutions, although being effective with respect to the problem of H₂O removal, rely on complex techniques and methodologies for carrying out them.

[0005] Other simpler technological solutions rely on a dispersion of active elements in polymers, in which the dimensions of the particles are utilized to obtain secondary properties, as the transparency, of the sorber produced in this way.

[0006] This type of solution is described in the patent US 6740145, where it is requested that the dimensions of the particles of the dispersed active element are between 1 and 100 nm, and in the patent application US 2006/0097633 where a average particle size in the polymeric film of less then 100 nm with a specific size distribution is requested.

[0007] The patent application MI2007/A000690, in the name of the applicant and not yet published on the filing

date of present application, shows the use of nanostructured fibres having inside an active component in form of dispersed particles.

[0008] The solutions shown in these documents have mainly two disadvantages, one inherent to the production and the other to the characteristics of the product. As relating to the disadvantage inherent to the production, this is associated to the use of particles of the active medium with a well defined particle size that requires also special precautions for their handling in case in which said particles have a very small diameter, i.e. 100 nm or less.

[0009] From the point of view of the efficiency of secondary characteristics of the product, i.e. not directly correlated to its H₂O sorbing capacity, as for example the transparency, the use of an active element in form of particles dispersed in a polymeric matrix can cause the deterioration in the course of time, mainly due to problems of particle aggregation inside the dispersion medium. This problem is increasingly relevant, the larger the particles sizes of the active medium are; the importance and criticality of the particle size of the hygroscopic material are pointed out several times in the description of the above-mentioned patent US 6740145. Object of the present invention is thus to overcome the inconveniences still present in the prior art as relating to the manufacturing of a H₂O sorber containing a hygroscopic inorganic salt, with special reference to the necessity to use inorganic salts of nanometric dimensions in which the particle size is of fundamental importance for the characteristics and properties of the produced sorber. In a first aspect thereof, the invention consists in a method for the manufacturing of a sorber for the removal of H₂O comprising a polymeric matrix and a hygroscopic inorganic salt, characterised in that said hygroscopic inorganic salt is dissolved inside of said polymeric matrix. In the following the term composite material will be used to identify the sorber material in order to highlight that is obtained dissolving an inorganic compound in a polymerized organic mixture, even if it is actually an homogeneous material that is not characterized by phase separation.

[0010] The invention will be illustrated in the following with reference to the drawings, in which:

- 45 - figure 1 shows a photography taken by an optical microscope of a composite sorber made according to the present invention;
- figure 2 shows a photography taken by an optical microscope of a composite sorber made with one of the inorganic salts suitable for carrying out the present invention, but not with the method herein described;
- figure 3 shows a photography taken by an optical microscope of a composite sorber made with a non-suitable material;
- figure 4 show the comparison of the transparency of different composite polymeric films; and
- figure 4A shows the comparison between the trans-

parency of a film, made according to the method of the invention, at time zero and of the same film after 23 hours of exposure to air.

[0011] Differently from the solutions of the prior art, for the manufacturing of the invention it is also possible to use powders with a particle size larger than 5 μm , and even, as will be illustrated in the following, also with a particle size larger than 2 mm; this simplifies their storage and handling, other than providing advantages in the manufacturing process, as will be illustrated further ahead.

[0012] The inventors have discovered various methods to manufacture the H_2O composite sorbers consisting of a polymeric matrix and a dissolved hygroscopic inorganic salt; in particular the polymer and the inorganic salt are both dissolved in a solvent, the evaporation of which results in the consolidation of the polymeric matrix that thus contains the dissolved inorganic salt. In a preferred embodiment the polymer and the inorganic salt are dissolved in the same common solvent, but it is also possible to use two different solvents for the polymer and for the hygroscopic inorganic salt, in which case it is necessary that the two solvents are miscible with each other.

[0013] In alternative it is possible to directly dissolve the hygroscopic inorganic salt in the polymer precursor, ideally when this is still in the monomer state. Subsequently the polymerization process, for example by thermal treatment or UV irradiation, leads to the formation of the polymeric matrix inside which the hygroscopic inorganic salt is dissolved.

[0014] The information on the solubility of polymers and inorganic salts in solvents, the miscibility of different solvents, the solubility of inorganic salts in polymer precursors (their monomers, that in this case act as solvent for the inorganic salt) are widely known and diffused and available to experts in the field and can be found, for example, in publications as the *Handbook of Chemistry and Physics*, 87th edition, 2006-2007, edited by CRC or *Alkaline Earth Metal Perchlorates*, IUPAC Solubility data Series, Vol. 41, edited by Pergamon Press. Solubility data can be moreover found in other publications, as for example the *Handbook of Solubility Parameters and other Cohesion Parameters*, 2nd edition, 1991, or *Hansen Solubility Parameters - A user's Handbook* by Charles M. Hansen, 2nd edition, 2007 both edited by CRC Press.

[0015] Transparent electrolytes for lithium batteries comprising a suitable polymer with an organic/inorganic compound, in form of gel, are described in the article "Ion-conductive polymethylmethacrylate gel electrolytes for lithium batteries", published in 2005 (available online 27 April 2005) on the *Journal of Power Sources*, 146, pages 436-440.

[0016] Whenever the solubility of a salt in a solvent, a monomer, a polymer or a mixture of at least two of them is not directly available in bibliographic data, an expert in the field can easily obtain it as experimental value. In fact a simple solubility test can consist in subsequent

addition of small amount of solute (salt) in a fixed amount of the liquid mixture: starting of precipitate formation correspond to the solubility limit of the salt in the chosen mixture.

[0017] Hygroscopic inorganic salts suitable for carrying out the invention are alkaline-metal and alkaline earth-metal perhalogenates, alkaline-metal and alkaline earth-metal halides; the use of perchlorates is preferred.

[0018] In the case of composite sorbers that foresee the use of solvents, some examples of polymers and solvents suitable for carrying out the invention are cellulose acetate in methyl acetate, or the same cellulose acetate in tetrahydrofuran cellulose.

[0019] In the case in which the solution is composed only of the polymer precursor and of the hygroscopic inorganic salt (the precursor acts as solvent for the inorganic salt) some examples of suitable polymers are polymethylmethacrylate (PMMA), obtained by polymerization of methylmethacrylate (MMA), or polyethylmethacrylate (PEMA) obtained by polymerization of ethyl methacrylate (EMA).

[0020] It is also possible to use combinations of polymers or polymer precursors for obtaining the polymeric matrix of the composite sorber being object of the present invention. Moreover, the polymeric matrix is formed starting from two different polymers, two precursors or a mixture of a polymer and a precursor. Furthermore said precursor can be a monomer. The case in which one of these polymers or polymeric precursors acts inside the solution as cross-linking agent results to be particularly advantageous. As an example, the use of ethylene glycol dimethylacrylate as cross-linking agent for obtaining a composite sorber having as polymeric matrix PMMA and as hygroscopic inorganic salt dissolved therein magnesium perchlorate results to be particularly advantageous. Moreover particularly advantageous is the combination of a polymeric precursor with a low molecular weight polymer. As an example, the use of mixture of polymethylmethacrylate (PMMA) with methylmethacrylate (MMA) results to be advantageous whenever the outgassing and shrinkage amount must to be minimized, i.e. if the polymerization has to take place in a closed environment or in a closed device.

[0021] Functionalized polymer precursors can moreover be added in order to improve and optimize physicochemical material properties. The adhesion on a selected substrate in fact is very important for the final application of a composite sorber and the use of trimethylsiloxyethyl methacrylate results to be very advantageous to improve the adhesion on a glass-type or a metal-oxidized substrate.

[0022] Other additives are usually foreseen in the final composition in order to act as polymerization promoter. Different kinds of polymerization catalysts can be used, as for example cationic, anionic or radical initiator, and it is chosen in function of the polymerization process that will be used to obtain the final matrix. Its concentration is usually lower or equal to 1 %w/w.

[0023] The quantity of inorganic salt to be added during the manufacturing process of the composite sorber depends on the specific hygroscopic inorganic salt used and on the characteristics of the polymer precursor (in the case in which this acts as solvent for the inorganic salt) or on the solvent-polymer-inorganic salt combinations and is difficult to determine a priori; and it is thus necessary to gradually add quantities of the inorganic salt to dissolve to the solution that will allow to obtain the polymeric composite sorber, proceeding with its dissolution, for example by stirring the solution and observing when the addition of the inorganic salt corresponds to the formation of a precipitate on the bottom of the solution. When this situation is reached this indicates that the addition of further inorganic salt quantities will not increase in any way the concentration of the dissolved inorganic salt, whereby it is possible to proceed to the consolidation of the polymeric matrix, upon eliminating the precipitated inorganic salt, for example by means of filtration.

[0024] The use of hygroscopic inorganic salt powders with a particle size larger than $5\mu\text{m}$ makes more evident the determination of the limit conditions being reached and simplifies the subsequent operations of removing the excess precipitate.

[0025] Obviously it is also possible that the saturation condition of the solution is not reached, in which case the removal operation of the precipitated inorganic salt is not necessary.

[0026] In a second aspect thereof the invention consists in a method for the removal of H_2O from devices sensitive to its presence by the use of composite sorbers consisting of hygroscopic inorganic salts dissolved in a polymeric matrix.

[0027] Among the sensitive devices that benefit the most from the application of the method of the invention are photovoltaic cells, OLED displays, microelectromechanical devices and lithium batteries.

[0028] In general the method of the invention offers advantages when it is necessary that the H_2O concentration inside the sensitive device must not exceed a critical value during the normal functioning of the device. This critical value is related to the kind of sensitive device and among those which require a very low water concentration are the OLEDs, that typically need concentrations in the order of 10 ppm or less, whereas at the extreme opposite there are the solar cells, which can support up to 5000 ppm before irreversible deterioration phenomena are triggered.

[0029] The polymer containing the dissolved hygroscopic inorganic salt may be used in form of an already consolidated film, in this case with thicknesses typically between $10\mu\text{m}$ and $200\mu\text{m}$, or it may be used when not yet completely solidified, while carrying out the final phase of consolidation or polymerisation once it has been dispensed: in this case the use of the monomer as medium in which the hygroscopic inorganic salt is dissolved results to be particularly advantageous for this way of

use, because there is no evaporation of the solvent that could give rise to contaminations of the device. The dispensing on the final support may be carried out by various methods widely known in the field, for example through brush work or spraying; preferred is the use of serigraphic method, well known in the printing field, which allows a better control of the deposit thickness (by means of control of the stencil thickness, through which the mixture is forced to pass for reaching the support) or also a filmograph may be used (a plate held at a fixed distance from a base or support, at a distance that corresponds to the thickness of the film). Moreover, said dispensation occurs by serigraphy, in which the final step of consolidation is a thermal or UV promoter polymerization.

[0030] Another way of using the sorbers made according to the present invention foresees the melting of the polymer containing the dissolved hygroscopic inorganic salt; in this case it is useful to obtain the H_2O sorber from a thermoplastic polymer, i.e. from a polymeric material with a melting temperature typically lower than 300°C .

[0031] Another embodiment foresees the use of H_2O sorbers made according to the present invention in the form of nanofibres, that can be produced by a technique known in the field by the term "electrospinning".

[0032] In some devices it may be useful to fill the inner volume of the sensitive device with the composite sorber; in this case the preferred solution foresees the introduction of the composite sorber in already consolidated form. A subsequent thermal treatment causes then the melting thereof and consequent filling of the inner volume of the device in a nearly uniform manner. Among the sensitive devices that benefit the most of this particular configuration are the photovoltaic cells and the OLED displays.

[0033] In a third aspect thereof the invention is inherent to composite sorbers consisting of polymeric matrices comprising dissolved hygroscopic inorganic salts that could be chosen from one or more of the following inorganic compounds: alkaline-metal and alkaline earth-metal perhalogenates, alkaline-metal and alkaline earth-metal halides, among which the use of perchlorates results to be preferred.

[0034] The invention will be further described with reference to the following examples:

45 **Example 1**

[0035] A film of a polymeric composite sorber according to the present invention is produced by dissolving a hygroscopic inorganic salt in a monomer. A quantity of 0.4 grams of magnesium perchlorate is used, the powder's particle size of which is not controlled and may also comprise flakes of a diameter of 2 mm. The inorganic salt is dissolved in 5 grams of methylmethacrylate (MMA) and 0.05 grams of benzoin methyl ether, this last one having the function of polymerisation initiator.

[0036] The polymerisation is carried out inside a chamber model UVACUBE 100 produced by Honle, and obtained by means of irradiation with ultraviolet light pro-

duced by a 100 W mercury lamp. In the following example, the solution is subjected to a "pre-UV curing" treatment for 14 minutes with a radiation dose equal to 4.35J/cm².

[0037] The pre-polymerised solution is spread out by means of a filmograph with a thickness of 50 microns on a steel plate and finally consolidated by means of a "post-UV curing" treatment for 30 minutes (the corresponding radiation dose being 9.32J/cm²). All of the preceding operations are carried out in a glove box under an inert gas flux in order not to compromise the sorbing capacity of the hygroscopic inorganic salt.

[0038] The photography by optical microscopy is shown in figure 1.

Example 2 (comparative)

[0039] A film of a polymeric composite sorber is produced by not operating under the conditions of the invention i.e. using a common solvent, dichloromethane (CH₂Cl₂), for the polymer (PMMA) and the inorganic salt (Mg(ClO₄)₂), in which the solvent is not able of dissolving the inorganic salt.

[0040] The quantities of polymer and inorganic salt are the same as in example 1, also the particle size of the inorganic salt is the same, whereas as regards the quantity of the solvent, a quantity equal to 15g has been used.

[0041] The photography by optical microscopy is shown in figure 2.

Example 3 (comparative)

[0042] A film of a polymeric composite sorber is produced by operating as in example 1, i.e. using a solution given by the monomer (MMA) mixed with powders of hygroscopic material, but using a hygroscopic material not foreseen by the present invention, i.e. calcium oxide in powder form, added in a quantity equal to 0.4 g, that does not dissolve in methylmethacrylate.

[0043] The photography by optical microscopy is shown in figure 3.

Example 4

[0044] This example compares a secondary characteristic, the transparency, of the films from polymeric composite sorbers obtainable with some special polymers (among which PMMA) prepared as described in the examples 1-3.

[0045] For this characterisation a double beam spectrophotometer Jasco V 570 has been used, with a single monochromator, a Hamamatsu detector, using deuterium and halogen lamp, scanning velocity 1000 nm/min, passband 1 nm and 0.3 nm accuracy. The characterised area of each polymeric film corresponds to the area of the incident light beam on the sample, i.e. a rectangle of 1 x 10 mm².

[0046] The results related to the spectral interval

400-700 nm are shown in the figures 4 and 4A in which:

- curve 1 shows the transmission curve of a PMMA film without any hygroscopic inorganic salt;
- the curves 2 and 2' show the transmission curves of a film prepared according to what described in example 1, respectively at time 0 and after 23 hours of exposure to air. In order to distinguish the two curves, which result to be almost perfectly superimposed, circles have been used to show the points for the test at time 0, whereas crosses have been used for the test after 23 hours; and
- curve 3 shows the transmission curve of a film prepared according to what described in example 2, i.e. a PMMA film comprising not perfectly dissolved magnesium perchlorate salts.

[0047] In figure 4A the comparison between the transparency curve at time 0 (curve 2a) and after 23 hours of exposure to air, (curve 2a'), is shown, using a very expanded scale for the axis of ordinates in order to appreciate the differences between these curves.

[0048] It was not possible to add a curve relating to the transparency of a sample prepared according to example 3, because this already at first sight resulted opaque.

[0049] Figures 1-3 highlight how only in the case of figure 1 there are no particles or their aggregates in the polymeric composite sorber film, as instead happens in the case of figure 2, in which there is not perfectly dissolved magnesium perchlorate in the film which gives rise to a film with irregular characteristics, whereas figure 3 shows the calcium oxide particles enclosed in the polymeric film.

[0050] Figure 4 allows to observe how the morphological differences in the polymeric composite sorber film can transform also in phenomenological differences, referring particularly to a secondary characteristic of the hygroscopic film, i.e. its transparency. This secondary characteristic is very useful in the case the sensitive device is an OLED display or placed in a solar cell.

[0051] In fact it can be observed how a hygroscopic film produced according to the present invention shows a transparency (lines 2, 2') that is totally comparable to that of a polymer without inorganic salt (line 1). Further the characteristic of transparency of the hygroscopic film is not compromised by the H₂O sorbing of the active component contained therein, i.e. the hygroscopic inorganic salt, and results to be always greater than 95% in the spectral interval considered.

[0052] Figure 4A shows how there are no significant differences in the transmission curve of a composite sorber obtained according to the method of the invention at time zero (curve 2a) and after 23 ours of exposition to air (curve 2a').

[0053] Instead a film containing a hygroscopic inorganic salt not perfectly dissolved in the polymeric matrix, apart from showing in general definitely inferior transmission characteristics (curve 3) with respect to the films of

the present invention, has also a high variability of its transparency characteristics according to the considered sample zone of the film.

[0054] The situation is still more critical if an active component is chosen to be added to the polymeric matrix that is not among those described in the present application, as in the case of calcium oxide, that results to give rise to an opaque hygroscopic film and for which it was not even possible to perform the characterisation.

Claims

- Method for manufacturing a composite sorber for the removal of H_2O comprising a polymeric matrix and a hygroscopic inorganic salt, **characterised in that** said hygroscopic inorganic salt is dissolved in a polymer precursor or a mixture of at least a polymer and a polymer precursor, that provides a solvent for the hygroscopic inorganic salt.
- Method according to claim 1, in which said hygroscopic inorganic salt is chosen from alkaline-metal and alkaline earth-metal perhalogenates, alkaline-metal and alkaline earth-metal halides.
- Method according to claim 2, in which said alkali metal and alkaline earth-metal perhalogenates are perchlorates.
- Method according to claim 1, in which said polymeric matrix is formed starting from two different polymers, two precursors or a mixture of a polymer and a precursor.
- Method according to claim 4, in which one of said polymers or of said polymer precursors acts as cross-linking agent for the polymeric matrix.
- Method according to claim 1, in which said sorber is manufactured by thermal treatment or UV irradiation from a solution comprising the polymer and the dissolved hygroscopic inorganic salt.
- Method according to claim 6, in which said solution comprises a common solvent for the polymer and the hygroscopic inorganic salt.
- Method according to claim 1, in which said polymer precursor is its monomer.
- Method for the removal of H_2O from devices sensitive to its presence by the use of composite sorbers comprising a hygroscopic inorganic salt dissolved in a polymeric matrix, **characterized in that** said hygroscopic inorganic salt is chosen from the group of alkali-metal and alkaline earth-metal perhalogenates, alkali-metal hydroxides, alkali-metal and alka-

line earth-metal halides and their combination.

- Method according to claim 9, in which said sensitive device is a OLED display.
- Method according to claim 9, in which said sensitive device is a microelectromechanical device.
- Method according to claim 9, in which said sensitive device is a solar cell.
- Method according to claim 9, in which said sensitive device is a lithium battery.
- Method according to claim 9, in which said composite sorber is used in form of a thin film with a thickness between 10 and 200 μm .
- Method according to claim 9, in which said composite sorber is dispensed inside of said sensitive device before the final step of consolidation of said sorber.
- Method according to claim 15, in which said dispensation occurs by serigraphy.
- Method according to claim 15, in which the final step of consolidation is a thermal or UV-promoted polymerization.
- Method according to claim 9, in which said polymeric composite sorber has the form of nanofibres.
- Method according to claim 18, in which said nanofibres are manufactured by the technique of electrospinning.
- Method according to claim 9, in which said sorber is introduced in said sensitive device by melting of said composite sorber.

Patentansprüche

- Verfahren zur Herstellung eines VerbundSORPTIONSMittels zur Entfernung von H_2O , umfassend eine Polymermatrix und ein hygrokopisches anorganisches Salz, **dadurch gekennzeichnet, dass** das hygrokopische anorganische Salz in einem Polymerprecursor oder einer Mischung von mindestens einem Polymer und einem Polymerprecursor, die das Lösungsmittel für das hygrokopische anorganische Salz bereitstellt, aufgelöst wird.
- Verfahren nach Anspruch 1, wobei das hygrokopische anorganische Salz ausgewählt ist aus Alkali-metall- und Erdalkalimetallperhalogenaten, Alkali-metall- und Erdalkalimetallhalogeniden.

3. Verfahren nach Anspruch 2, wobei die Alkalimetall- und Erdalkalimetallperhalogenate Perchlorate sind.

4. Verfahren nach Anspruch 1, wobei die Polymermatrix ausgehend von zwei verschiedenen Polymeren, zwei Precursoren oder einer Mischung eines Polymeren und eines Precursors gebildet wird.

5. Verfahren nach Anspruch 4, wobei eines der Polymeren oder der Polymerprecursor als Vernetzungsmittel für die Polymermatrix fungiert.

6. Verfahren nach Anspruch 1, wobei das Sorptionsmittel durch thermische Behandlung oder UV-Be- strahlung einer Lösung, umfassend das Polymer und das gelöste hygrokopische anorganische Salz, hergestellt wird.

7. Verfahren nach Anspruch 6, wobei die Lösung ein übliches Lösungsmittel für das Polymer und das hygrokopische anorganische Salz umfasst.

8. Verfahren nach Anspruch 1, wobei der Polymerpre- cursor sein Monomer ist.

9. Verfahren zur Entfernung von H_2O aus Geräten, die sensibel auf dessen Präsenz sind, durch Verwen- dung von Verbundsorptionsmitteln, umfassend ein hygrokopisches anorganisches Salz, aufgelöst in einer Polymermatrix, **dadurch gekennzeichnet, dass** das hygrokopische anorganische Salz aus- gewählt ist, aus der Gruppe, bestehend aus Alkalimetall- und Erdalkalimetallperhalogenaten, Alkalimetall-hydroxiden, Alkalimetall- und Erdalkalimetall- halogeniden und deren Mischungen.

10. Verfahren nach Anspruch 9, wobei das empfindliche Gerät ein OLED Display ist.

11. Verfahren nach Anspruch 9, wobei das empfindliche Gerät ein mikroelektromechanisches Gerät ist.

12. Verfahren nach Anspruch 9, wobei das empfindliche Gerät eine Solarzelle ist.

13. Verfahren nach Anspruch 9, wobei das empfindliche Gerät eine Lithiumbatterie ist.

14. Verfahren nach Anspruch 9, wobei das Verbund- sorptionsmittel in Form eines dünnen Films mit einer Dicke zwischen 10 und 200 μm verwendet wird.

15. Verfahren nach Anspruch 9, wobei das Verbund- sorptionsmittel im Inneren des empfindlichen Gerätes verteilt wird, bevor der finale Schritt der Festigung dieses Sorptionsmittels erfolgt.

16. Verfahren nach Anspruch 15, wobei die Verteilung

5 durch Siebdruck erfolgt.

17. Verfahren nach Anspruch 15, wobei der finale Schritt der Festigung eine thermische oder UV-geförderte Polymerisation ist.

10 18. Verfahren nach Anspruch 9, wobei das polymere Verbundsorptionsmittel die Form von Nanofasern hat.

19. Verfahren nach Anspruch 18, wobei die Nanofasern durch Elektrospinnen hergestellt werden.

15 20. Verfahren nach Anspruch 9, wobei das Zuführen des Sorptionsmittels in das empfindliche Gerät durch Schmelzen des Verbundsorptionsmittels erfolgt.

Revendications

1. Procédé de fabrication d'un sorbant composite pour l'élimination de H_2O comprenant une matrice polymère et un sel inorganique hygroscopique, **carac- térisé en ce que** ledit sel inorganique hygroscopique est dissous dans un précurseur de polymère ou dans un mélange constitué d'au moins un polymère et d'un précurseur de polymère, qui constitue un solvant pour le sel inorganique hygroscopique.
2. Procédé selon la revendication 1, dans lequel ledit sel inorganique hygroscopique est choisi parmi les perhalogénates de métal alcalin et de métal alcalino-terreux, et les halogénures de métal alcalin et de métal alcalino-terreux.
3. Procédé selon la revendication 2, dans lequel lesdits perhalogénates de métal alcalin et de métal alcalino-terreux sont des perchlorates.
4. Procédé selon la revendication 1, dans lequel ladite matrice polymère est formée en partant de deux polymères différents, deux précurseurs ou un mélange constitué d'un polymère et d'un précurseur.
5. Procédé selon la revendication 4, dans lequel un desdits polymères ou desdits précurseurs de polymères agit comme un agent de réticulation pour la matrice polymère.
6. Procédé selon la revendication 1, dans lequel ledit sorbant est fabriqué par traitement thermique ou ex- position à des UV à partir d'une solution comprenant le polymère et le sel inorganique hygroscopique dis- sous.
7. Procédé selon la revendication 6, dans lequel ladite solution comprend un solvant commun pour le poly- mère et le sel inorganique hygroscopique.

8. Procédé selon la revendication 1, dans lequel ledit précurseur de polymère est son monomère.
9. Procédé d'élimination de H₂O des dispositifs sensibles à sa présence par utilisation de sorbants composites comprenant un sel inorganique hygroscopique dissous dans une matrice polymère, **caractérisé en ce que** ledit sel inorganique hygroscopique est choisi dans le groupe des perhalogénates de métal alcalin et de métal alcalinoterreux, des hydroxydes de métal alcalin, des halogénures de métal alcalin et de métal alcalinoterreux et leur combinaison. 5
10. Procédé selon la revendication 9, dans lequel ledit dispositif sensible est un affichage OLED. 10 15
11. Procédé selon la revendication 9, dans lequel ledit dispositif sensible est un dispositif microélectromécanique. 20
12. Procédé selon la revendication 9, dans lequel ledit dispositif sensible est une cellule solaire.
13. Procédé selon la revendication 9, dans lequel ledit dispositif sensible est une batterie au lithium. 25
14. Procédé selon la revendication 9, dans lequel ledit sorbant composite est utilisé sous la forme d'un film mince ayant une épaisseur comprise entre 10 et 200 µm. 30
15. Procédé selon la revendication 9, dans lequel ledit sorbant composite est distribué à l'intérieur dudit dispositif sensible avant l'étape finale de consolidation dudit sorbant. 35
16. Procédé selon la revendication 15, dans lequel ladite distribution s'effectue par sérigraphie.
17. Procédé selon la revendication 15, dans lequel l'étape finale de consolidation est une polymérisation thermique ou UV. 40
18. Procédé selon la revendication 9, dans lequel ledit sorbant composite polymère est sous la forme de nanofibres. 45
19. Procédé selon la revendication 18, dans lequel lesdites nanofibres sont fabriquées par la technique d'électrofilage. 50
20. Procédé selon la revendication 9, dans lequel ledit sorbant est introduit dans ledit dispositif sensible par fusion dudit sorbant composite.

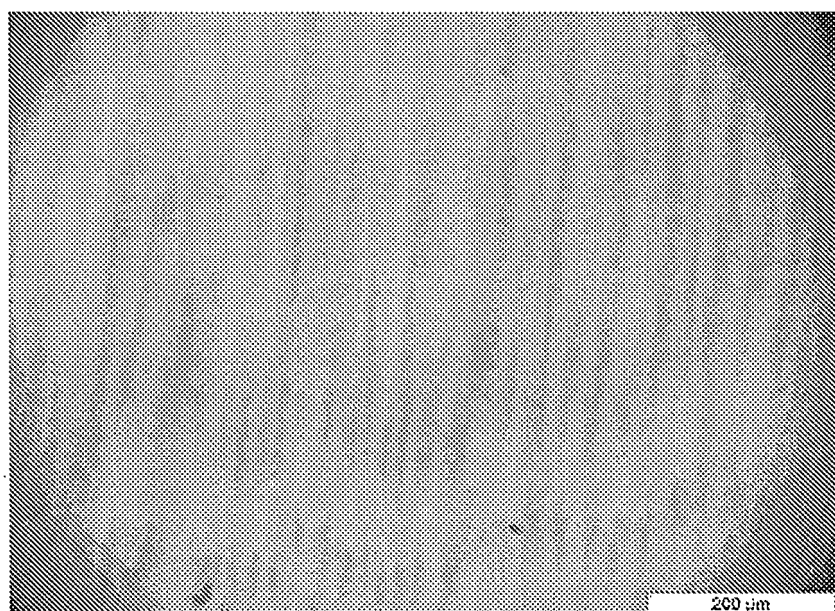


Fig. 1



Fig. 2

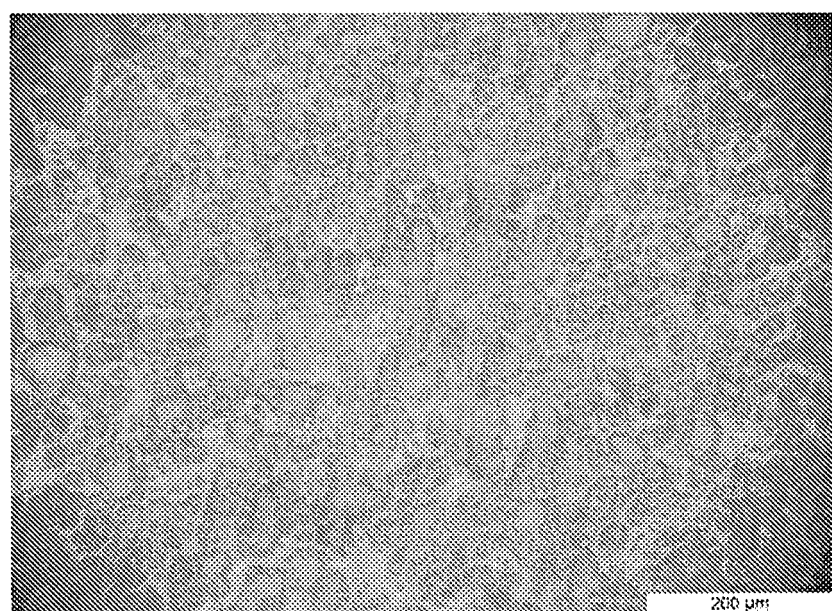


FIG. 3

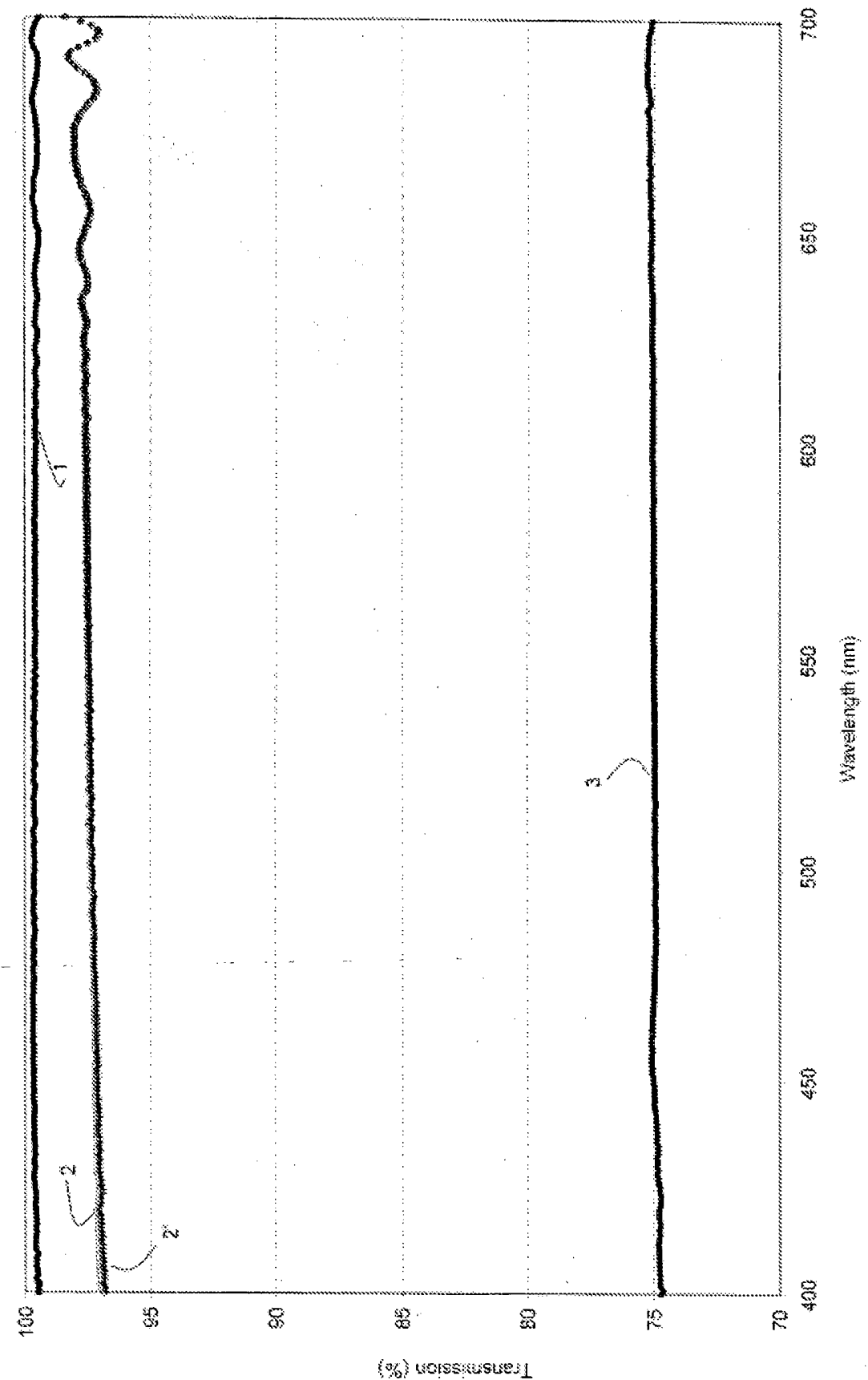


FIG. 4

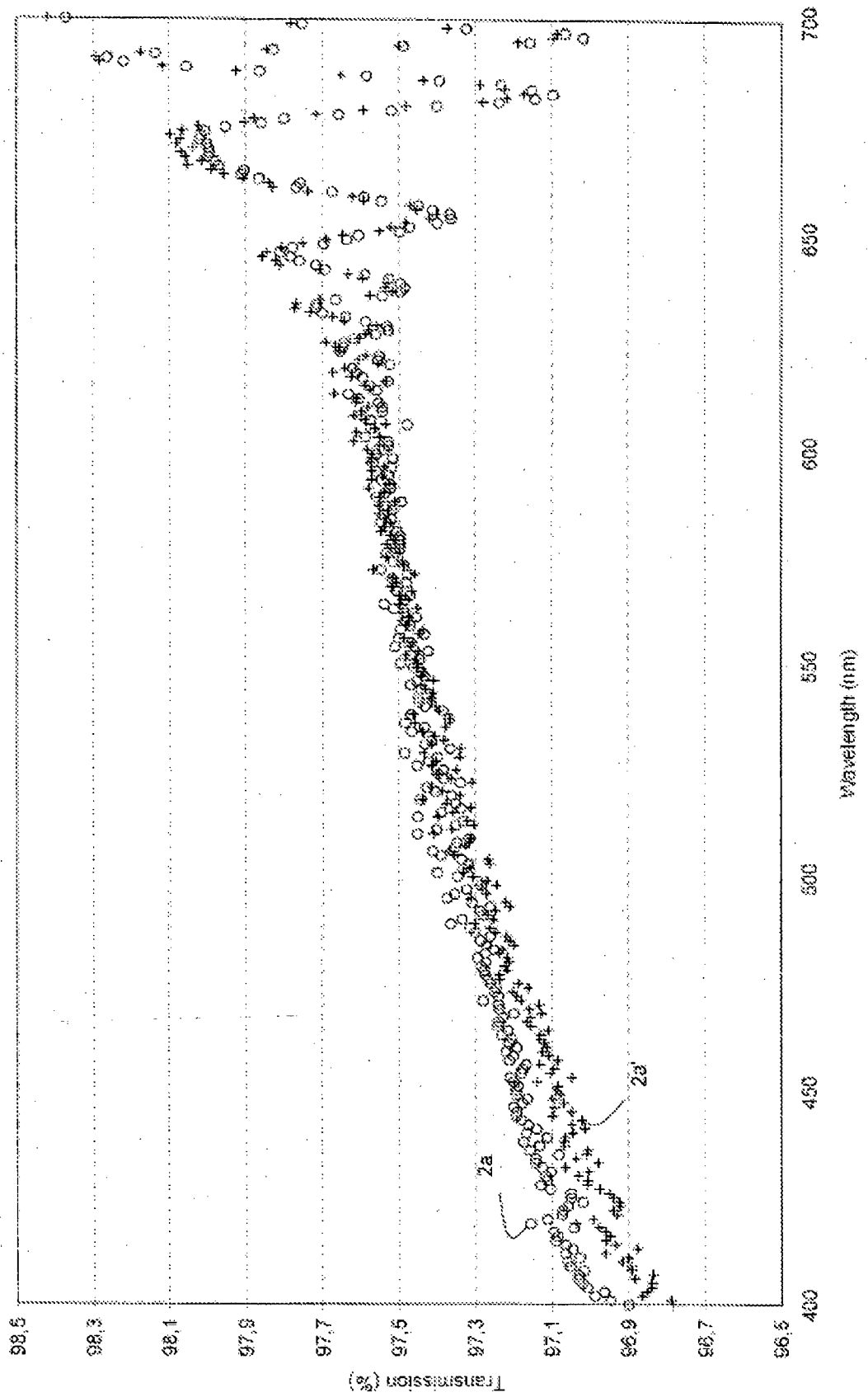


FIG. 4A

REFERENCES CITED IN THE DESCRIPTION

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